

PATENT TS0633 (US) DFH:EM

### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of

JOCHEM VAN DE WEERD

Serial No. 10/808,001

Filed March 24, 2004

METHOD OF INCREASING THE CARBON CHAIN )
LENGTH OF OLEFINIC COMPOUNDS )

May 11, 2007

COMMISSIONER FOR PATENTS P. O. Box 1450 Alexandria, VA 22313-1450

Madam/Sir:

#### DECLARATION UNDER RULE 131

Dr. ARIE VAN ZON hereby déclares:

THAT he is a Principal Research Chemist for Shell Global Solutions International

BV; and

THAT he has reviewed WO 03/024910 and understands that this patent application is one of the references used by the Examiner in the office action of March 27, 2007 to reject the claims of the present application; and that the priority date of this reference is September 17, 2001; and

THAT he is familiar with the request for patent action and the report describing the work which was done relating to the present invention; and that the request for patent action and the work referred to above were written and performed in the Netherlands, a WTO country, prior to September 17, 2001; and

THAT the document entitled "Request for Patent Action" enclosed herewith was created prior to September 17, 2001 and describes the invention which is claimed in the present application; and that the document entitled "Dehydration of SHF-Alcohols to Alpha Olefins" was created prior to September 17, 2001; and that this document describes experimental work wherein a primary alcohol, i.e. pure (99%) 1-decanol, and industrially produced primary alcohol mixtures, such as NEODOL-1 (which is largely linear 1-undecanol) or Linevol-911 (a mixture of

 $C_{\theta}$ ,  $C_{10}$  and  $C_{11}$  1-alcohols), made by hydroformylation of a  $C_{10}$  olefin or a mixture of  $C_{\theta}$ ,  $C_{\theta}$  and  $C_{10}$  olefins were subjected to dehydration in a continuous reactor in the gas phase and that 1-decene, 1-undecene or a mixture of 1-nonene, 1-decene and 1-undecene, respectively, were the predominant products; and

THAT the attached documents show that the invention claimed in the present application was made prior to September 17, 2001.

Dr. Arie van Zon further declares that all statements herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Date 11th May 2007

CONFIDENTIAL

To be sent to: IP/43

The Hague

Q be completed by Intellectual Property Division

0633

Class(es): 15405 Field Code: CH rt Title: PROCESS FOR CONVERTING Cx TO Cx+1 OLEFINS

Date Received:

Attorney: LAVY

To be completed by Reporter

Invention originating from: SRTCA

Department: CTHOC

Research Budget Code(s): (Code with format \*\*\*.\*\*.\*\*) 153.151.01

Brief Description of Research A process for converting a first olefin having x carbon atoms to a second olefin having x+1 carbon atoms, which process comprises:

OUEST FOR PATENT ACTION

a hydroformylation stage wherein the first olefin is reacted with carbon monoxide and hydrogen in the presence of a hydroformylating catalyst to form the corresponding alcohol. followed by

a dehydration stage wherein the alcohol is dehydrated in the presence of a dehydration catalyst to form the second olefin.

The process is of special importance in the conversion of low-value olefins such as unevennumbered olefins originating from the Fischer-Tropsch reaction (especially 1-pentene) or from the SHOP oligomerisation (especially iso-nonene) to the corresponding higher-valued even-numbered alpha olefins. Preferred hydroformylation and dehydration catalysts guarantee selectivity towards alpha olefinic end products.

Name(s) of Contributor(s): J. M. van de Weerd

Are there any relevant reports and/or experimental evidence available? (If YES, please attach copies

NO

Are there any relevant patents or other literature known? (If YES, please attach details)

Name of Reporter U. Lavy Extension No.: 2095 Date:

Abundant literature about each of the two reactions Co-SHF is a selective first step and GB-A 2181070 (K605) discloses a selective

second step. Signature of Departme

From: TABS Editorial Office, CTAIC2, SRTCA, tel (+31)20 630 2344
Subject: CONF: TABS.

TITLE: Dehydration of SHF-alcohols to alpha olefins

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AUTHOR: N. Meijboom (SRTCA-CTHOD/1)

KEYWORDS: dehydration, decanol, decene, NEODOL, PAO

### INCENTIVE:

Preliminary desk studies have been carried out to evaluate the possibilities of combining a Fischer-Tropsch (FT) unit with a higher olefins and derivatives (HODER) and lower olefins (LO) unit (1) A recent study (1a) is based largely on proven technology and its outcome was promising. The HODER, part of the scheme consists of a series of hydroformylation plants (SHP), which convert (internal) olefins to primary alcohols (NEODOL), followed by a dehydration unit, which converts the primary alcohol to an alpha-olefin. This dehydration step in the FTSHP units has not been entirely demonstrated and more detailed information is required.

# TECHNICAL SCOPE:

We carried out experimental scouting work on the dehydration of 1decanol, NEODOL-1 and Linevol-911. In collaboration with CTPUC/3

(2) we used an experimental setup designed to the simulate the unit in Moerdijk for MPC (methyl phenyl carbinol) dehydration to styrene. The dehydration was carried out in a continuous reactor in the gas phase (WHSV of 1240 kg.m-3.h-1) using the commercial MPC-dehydration catalyst(alumintum oxide ex-Engelhard, AL-0104- T). Well known side reactions in this chemistry are ether formation, olefin isomerisation and oligomerisation(3). In order to find the optimum dehydration conditions the initial experiments were carried out with pure (99%) 1-decanol, the results of which are summarised in table 1.

Table 1: Dehydration of 1-decanol to 1-decene (based on GLC-Area%)

vity	
selectivity to 1-decene 97.4% 96.5%	93.1%
didecyl total ther decenes 26.7% 51.4% 12.8% 73.6% 1.9% 92.5%	97.3%
n didecyl tr ether decenes 26.7% 51.49 12.8% 73.69 1.9% 92.5%	0.2%
cemp. Conversion oC) (%) ethe 26.82 79.3% 26.01 87.6% 12.13 96.2% 1.3	99.3%
Temp. (oC) 282 301 313	325

As can be seen in table 1 the best results (high conversion, low ether formation) were obtained at a temperature of 325 oC. At this temperature >99% of the alcohol is converted, the yield of decenes

is 97% and the selectivity to 1-decene is 93%. The two most abundant other decenes were tentatively identified as cis-2-decene (4.1%) and trans-2-decene (1.6%). It may be possible to leave the minor amounts of residual alcohol, ether and isomeric decenes in the product, as they are not expected to hamper the commercially used BF3/ROH catalysed oligomerisation of 1-decene to PAO (Poly Alpha Olefins).

These results led us to apply the same conditions to the dehydration of the SHF-alcohols Neodol-1 and Linevol-911. The dehydration of the SHF-alcohols Neodol-1 and Linevol-911. The dehydration of these alcohols proved to proceed just as smoothly as observed for 1-decanol at 325 oC. Conversion was again some 99% and ether formation as low as 0.2%. We will test the oleffins produced from C9-C11 range SHF alcohols for PAO application, because even in todays market situation the price difference between Linevol 911 and 1-decene is such that a low capex transformation like dehydration may be economically feasible from current commercial product (4). The dehydration of Neodol-1 gave a mixture of undecense with an 1-undecene content of 77.0%. Given the fact that the starting material Neodol-1 contained 83.5% 1-undecanol, the selectivity is comparable (92-93%) to the experiment with 1-decanol. The 2-alky branched alcohols (ca. 15-20%) in the Neodol give rise to a different set of alpha-olefins: vinylidene olefins.

Considering the increase in C-number from CIO to CII, the presence of these virylidene olefins could have a positive effect on PAO-properties. The dehydration of Linevol-911, with a normality (1-alcohol contem) of some 82% and containing ca. 19% C9, 45% CIO and 36% CII, gave a mixture of nonenes, decenes and undecenes. The mixture contained about 1% of starting material, 0.5% of ethers, about 15% of 1-nonene, 34% of 1-decene and 27% of 1-undecene. A total of about 76% 1-alkenes (the remainder being predominantly virylidene olefins) means again a selectivity of 92.5%, as the starting material contained only 82% of 1-alcohols. PAO-production and evaluation, using the above described olefins, will be subject of a future 7ABS.

In conclusion it can be stated that the dehydration of higher alcohols (C9-C11) to higher olefins proceeds smoothly with the MPC dehydration catalyst in the chosen equipment at 325 oC, showing high conversion (>92%). Although at lower temperatures the conversion and ether formation become unfavourable, the selectivity to alpha-olefins formation become unfavourable, the selectivity to 1 deceme is better (some 97%). A higher conversion at a lower temperature might be obtained with a modified alumina catalyst (5). The planning of further work depends on the results of the performance evaluation of PAO, derived from these olefins and the overall business

attractiveness of the Fischer-Tropsch based HODER-feedstock generator.

## REFERENCES:

1. a. H.Donkers, J. Ansorge, M. Crujisberg, J. van der Sluis; Gas to Chemiculs; CTBCH2623997 b. A. van Zun, H.J. van Milligen and A.K. van Helden; Review of the options for a HODER feedstock generator; PRC/036995 c. M.I. Barr, Cnecking of Bintlut pariffins, SICM-CMFP212 Note of 11 March 1993

Equipment was kindly made available to us by S.H. Johnson and J.C. van Reermund. The practical assistance of M. van Zwienen is gratefully

 R. Askani in Houben-Weyl, Methoden der Organischen Chemie, Band V/1B, viene auflage, 45-62; Eugen Moeller Toebingen (1972)

acknowledged.

Zinc aluminates are known dehydration catalysis that work at a lower temperature (e.g. C.S. John; Shell patent GB2181070A)

Communication with S.R. Brown (CED/11) d.d. 6-6-1997 5.

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